

# Proteomic Response of *Bacillus subtilis* to Lantibiotics Reflects Differences in Interaction with the Cytoplasmic Membrane

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Mersacidin, gallidermin, and nisin are lantibiotics, antimicrobial peptides containing lanthionine. They show potent antibacterial activity. All three interfere with cell wall biosynthesis by binding lipid II, but they display different levels of interaction with the cytoplasmic membrane. On one end of the spectrum, mersacidin interferes with cell wall biosynthesis by binding lipid II without integrating into bacterial membranes. On the other end of the spectrum, nisin readily integrates into membranes, where it forms large pores. It destroys the membrane potential and causes leakage of nutrients and ions. Gallidermin, in an intermediate position, also readily integrates into membranes. However, pore formation occurs only in some bacteria and depends on membrane composition. In this study, we investigated the impact of nisin, gallidermin, and mersacidin on cell wall integrity, membrane pore formation, and membrane depolarization in *Bacillus subtilis*. The impact of the lantibiotics on the cell envelope was correlated to the proteomic response they elicit in *B. subtilis*. By drawing on a proteomic response library, including other envelope-targeting antibiotics such as bacitracin, vancomycin, gramicidin S, or valinomycin, YtrE could be identified as the most reliable marker protein for interfering with membrane-bound steps of cell wall biosynthesis. NadE and PspA were identified as markers for antibiotics interacting with the cytoplasmic membrane.

ver the last decades, bacteria have demonstrated their impressive ability to adapt to changing environmental conditions by rapidly developing and accumulating antibiotic resistances. Helped by an extensive use of antibiotics in health care and agriculture, multidrug-resistant strains, so-called superbugs, are spreading with a remarkable impact on human health. Recent statistics of the World Health Organization show that even in developed countries, infectious diseases are back among the top five causes of death, with the majority of deaths being attributed to bacterial infections (49). At the same time, approval rates of new antibiotic agents have been steadily decreasing since their apex in the 1980s (5). In view of these developments, in addition to improving hospital hygiene and to devising and implementing strategies to preserve effectiveness of available antimicrobial agents, a reinvigoration of antibiotic research and development is urgently needed to meet the superbug challenge.

One of the promising antibiotic classes for treatment of infections caused by multidrug-resistant pathogens are antimicrobial peptides, which occur naturally as components of the host defense and typically target the bacterial cell envelope. The peptide subclass of lantibiotics, which are produced by Gram-positive bacteria, is characterized by containing the nonproteinogenic amino acid lanthionine and often also other unusual amino acids. However, the group of lantibiotics is structurally and mechanistically diverse and can be further divided into class A, comprising stretched, flexible peptides, and class B, with more globular structures. The majority of peptides in both classes have been shown to inhibit cell wall biosynthesis. In contrast to class B members, some class A lantibiotics are also capable of integrating into the bacterial membrane and of pore formation, a process facilitated by binding of cell wall precursor lipid II (7, 13). This mechanistic duality depicts an interesting advantage of lantibiotics over conventional single-mechanism antibiotics. It was shown for various antibacterial compounds that inhibit targets encoded by multiple genes that resistance development is much slower than for antibiotics inhibiting a single target of a specific metabolic pathway (11, 32). As two essential complex targets, the cell wall biosynthesis machinery and the cell membrane, need to be altered to prevent cellular damage by dual-mechanism lantibiotics, resistance development by target mutation and stress adaptation is effectively deferred (10, 32). Such properties highlight antimicrobial peptides as attractive antibacterial agents.

Here, we investigate three different lantibiotics. Nisin, the first lantibiotic ever described, belongs to class A. The bactericidal effects of nisin involve two distinct mechanisms. At lower concentrations, nisin forms aggregates with lipid II, thereby preventing them from participating in cell wall biosynthesis (23, 35). After reaching a certain concentration threshold, complexes of 8 nisin and 4 lipid II molecules are formed that integrate into the bacterial membrane. They form pores large enough for ions and small amino acids to pass through (9, 23). Structurally similar gallidermin also belongs to class A and inhibits cell wall biosynthesis by lipid II binding. Potassium leakage indicating pore formation was observed in some but not all bacteria (8); however, *B. subtilis* was not tested. Christ et al. were able to correlate pore formation with

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membrane thickness and fatty acid branching, suggesting that this second mechanism of action of gallidermin is dependent on membrane composition (16). In contrast to nisin, which needs to dock to lipid II for membrane integration and pore formation, gallidermin shows very high association and low dissociation constants for binding phospholipid bilayers, indicating that it readily integrates into the membrane independently of lipid II binding. The insertion of gallidermin into the bilayer already influences membrane properties without a necessity for pore formation (16). Class B lantibiotic mersacidin is structurally and mechanistically different from nisin or gallidermin. More globular in structure, it is not able to integrate into the membrane or to form pores, so its antibacterial activity is based solely on inhibition of cell wall biosynthesis by binding to lipid II. In contrast to vancomycin, a glycopeptide which does not integrate into the membrane either, mersacidin does not seem to bind the amino acid tail of lipid II. Rather, it binds to the disaccharide headgroup of the lipid II molecule and additionally interacts with the pyrophosphate, suggesting that lipid II-bound lantibiotic molecules are localized near the outer layer of the cell membrane (6, 14).

These three lantibiotics were chosen to investigate the proteomic response of B. subtilis to cell wall biosynthesis inhibition by lipid II binding coupled with different levels of interference with membrane integrity. The physiological response of B. subtilis manifest in the proteome after antibiotic stress was previously shown to correlate with the antibacterial mechanism (2, 4, 47) and contributed to elucidating the antibacterial mechanisms of novel antibiotic compounds (12, 48). Proteomic profiles of B. subtilis treated with several cell wall biosynthesis inhibitors have been reported (4, 39, 46), but proteomic signatures indicative of specific aspects of cell wall biosynthesis inhibition have not yet been described. Utilizing lantibiotics, which range in mechanism from just binding lipid II to binding lipid II and forming large pores, and by further drawing on an extensive library of proteomic response profiles previously described (4), we identified responder proteins indicative of the different lantibiotic mechanisms. To directly correlate the proteome profiles with the influence of mersacidin, gallidermin, and nisin on B. subtilis, the impact of the lantibiotics on membrane integrity was studied in this model organism. Lantibiotic interference with cell wall biosynthesis was examined microscopically, looking at cell shape after methanolacetic acid fixation. Potassium efflux measurements served to investigate pore formation. And finally, the influence of lantibiotics on the membrane potential was studied microscopically with the help of a green fluorescent protein (GFP)-MinD fusion protein. MinD is a cell division protein requiring an intact membrane potential for localization at the cell poles and the cell division plane (41). Based on the correlation of these physiological experiments with the proteomic response, marker proteins for different mechanisms of membrane interference could be proposed.

#### MATERIALS AND METHODS

Antibiotics. All antibiotic stock solutions were prepared at 10 mg/ml. Nisin was dissolved in 0.01 M HCl, gallidermin in double-distilled water (both were purified according to Bonelli et al. [8]), and mersacidin (Sanofi-Aventis, Paris, France) in methanol. Valinomycin and gramicidin A were dissolved in dimethyl sulfoxide (DMSO), and bacitracin was dissolved in double-distilled water (all from Sigma-Aldrich, St. Louis, MO). Vancomycin (Applichem, Darmstadt, Germany) was dissolved in DMSO. Gramicidin S was synthetized and purified according to Wadhwani et al. (45) and dissolved in DMSO. **Bacterial strains and growth conditions.** *Bacillus subtilis* 168 (trpC2) (1) was grown at 37°C under steady agitation in Belitzky minimal medium (BMM) (42). MICs were determined in a modified MIC test described previously to match the growth conditions of the proteome experiment, specifically using chemically defined medium and supplying sufficient oxygen (48). Briefly, in a test tube, 2 ml of defined medium was inoculated with  $5 \times 10^5$  bacteria per ml and incubated with different lantibiotic concentrations at 37°C under steady agitation for 18 h. The MIC was defined as the lowest concentration inhibiting visible growth. In growth experiments, bacterial cultures were treated with different antibiotic concentrations in mid-exponential phase after reaching an optical density at 500 nm (OD<sub>500</sub>) of 0.35. For physiological stress experiments, including proteomic studies, antibiotic concentrations were chosen that reduced growth rates to approximately 50 to 70% compared to that of the untreated control culture.

**Light microscopy.** *B. subtilis* 168 cultures were grown in BMM to an  $OD_{500}$  of 0.35 and subsequently treated with 0.75 µg/ml nisin, 6 µg/ml gallidermin, 30 µg/ml mersacidin, 10 µg/ml valinomycin, 0.025 µg/ml gramicidin A, 1.5 µg/ml vancomycin, 12 µg/ml bacitracin, or 1 µg/ml gramicidin S. After 15 min of antibiotic exposure, 200 µl of bacterial culture were immediately fixed in 1 ml of a 1:3 mixture of acetic acid and methanol. Five microliters of fixed cells were immobilized in 5 µl BMM containing 0.5% low-melting agarose at 40°C. Cells were observed with an Olympus BX51 microscope using a U-UCD8 condenser and a UPlanSApo 100XO objective. Pictures were taken using a CC12 digital color camera and cell imaging software (all components by Olympus, Hamburg, Germany).

**GFP-MinD localization.** *B. subtilis* 1981 GFP-MinD (41) was grown overnight in BMM. Cells were then inoculated to an  $OD_{500}$  of 0.1 in modified BMM containing xylose instead of glucose to induce expression of the GFP-MinD fusion protein. After reaching an  $OD_{500}$  of 0.35, cells were exposed to antibiotics at concentrations described above. After 15 min of antibiotic stress, 5  $\mu$ l of the bacterial culture was withdrawn and the nonfixed, nonimmobilized samples were imaged immediately in fluorescent mode using the described equipment with a U-LH100HGAPO burner and a U-RFL-T power supply (Olympus, Hamburg, Germany).

Potassium release. Potassium efflux experiments were performed using a microprocessor pH meter (pH 213; Hanna Instruments, Kehl, Germany) with an MI-442 potassium electrode and MI-409F reference electrode. The electrodes were preconditioned by immersing both the potassium selective and the reference electrodes in choline buffer (300 mM choline chloride, 30 mM MES, 20 mM Tris, pH 6.5) for at least 1 h before starting calibration or measurements. Calibration was carried out prior to each determination by immersing the electrodes in fresh standard solutions containing 0.01, 0.1, or 1 mM KCl in choline buffer. B. subtilis 168 was grown in BMM and harvested at an OD\_{500} of 1.0 to 1.5 (3,300  $\times$ g, 4°C, 3 min), washed with 50 ml cold choline buffer, and resuspended in the same buffer to an  $OD_{500}$  of 30. The concentrated cell suspension was kept on ice and used within 30 min. For each measurement, the cells were diluted in choline buffer (25°C) to an OD<sub>500</sub> of about 3. Antibiotics were applied at the concentrations described above. Calculations of potassium efflux were performed according to the equations established by Orlov et al. (31). Antibiotic-induced leakage was monitored for 3 min with readings taken every 10 s and expressed relative to the total potassium release induced by nisin.

**Proteome analysis.** Protein extracts for 2D-PAGE were prepared as described previously (48). In short, 5 ml of a bacterial culture in early exponential growth phase were exposed to 6  $\mu$ g/ml gallidermin, 30  $\mu$ g/ml mersacidin, 10  $\mu$ g/ml valinomycin, or 1  $\mu$ g/ml gramicidin S or left untreated as the control. When *B. subtilis* cultures were treated during exponential growth, nisin exhibited a very narrow window of concentrations that solicit a stress response without causing massive cell lysis. Addressing this problem, concentrations from 0.25 to 1.25  $\mu$ g/ml of nisin were used in parallel for each of the replicate labeling experiments. For the proteomic analysis, only those cultures were processed that showed growth

inhibition but no extensive cell lysis. Cultures treated with 0.75  $\mu$ g/ml (replicates one and two) and a culture treated with 0.5  $\mu$ g/ml nisin (replicate three) were chosen for proteome analysis based on the impact of nisin on the growth rate. After 10 min of antibiotic stress, proteins were pulse labeled with [<sup>35</sup>S]methionine for 5 min. Incorporation of radioactive methionine was stopped by adding 1 mg/ml chloramphenicol and an excess of cold methionine. For nonradioactive preparative gels, cells were treated with antibiotics for 30 min to allow sufficient protein accumulation for mass spectrometry. Cells were harvested by centrifugation, washed three times with Tris buffer, and disrupted by ultrasonication.

Two-dimensional (2D) gels were prepared as described previously (48). In short, 55  $\mu$ g of protein for analytical and 300  $\mu$ g for preparative gels were loaded onto 24-cm immobilized pH gradient (IPG) strips, pH 4 to 7 (GE Healthcare), by passive rehydration for 18 h followed by isoelectric focusing. After reduction and alkylation, proteins were separated in a second dimension by SDS-PAGE using 12.5% acrylamide gels. Images were analyzed as described by Bandow et al. (3) using Decodon Delta 2D 4.1 image analysis software (Decodon, Greifswald, Germany). Proteins found to be induced more than 2-fold in three independently performed biological experiments were defined as marker proteins.

Protein spots were excised from preparative 2D gels and transferred into 96-well microtiter plates. Tryptic in-gel digest and subsequent spotting of extracted peptides with matrix on a matrix-assisted laser desorption ionization (MALDI) target was carried out automatically with the Ettan spot handling workstation (Amersham Biosciences, Uppsala, Sweden) as described by Eymann et al. (20). MALDI-time of flight (TOF) as well as MALDI-TOF/TOF (consecutive TOF analysis) measurements were carried out on a 4800 MALDI-TOF/TOF analyzer (Applied Biosystems, Foster City, CA). For database searches, the Mascot search engine 2.1.04 (Matrix Science Ltd., London, United Kingdom) with a specific *B. subtilis* sequence database was used as described previously (48).

#### RESULTS

In order to obtain specific marker proteins that allow delineation of different mechanisms of action affecting the cell envelope, we investigated a set of lantibiotics, which inhibit bacterial cell wall biosynthesis while exhibiting different degrees of interference with the bacterial membrane. Mersacidin binds to lipid II without integrating into the cytoplasmic membrane, gallidermin binds to lipid II and can disrupt the structure of the membrane by insertion, and nisin binds to lipid II, integrates into the membrane, and leads to pore formation.

To find appropriate conditions for proteome analyses and the characterization of the effects of lantibiotics on the cell envelope, MICs of the lantibiotics were determined under conditions resembling those of the proteome experiment, using the chemically defined medium and incubation under steady agitation. The MICs were 30 µg/ml for mersacidin, 6 µg/ml for gallidermin, and 5 µg/ml for nisin. Based on these MICs, growth experiments were performed to identify antibiotic concentrations that reduce growth rates without causing massive cell lysis or completely stopping bacterial metabolism. This is essential, as proteomic experiments require a stress level high enough to induce a proteomic response but at the same time allowing protein biosynthesis to continue. While for mersacidin and gallidermin the MICs were optimal stressor concentrations, nisin caused massive cell lysis when applied at concentrations near the MIC during exponential growth. However, cells surviving the acute nisin shock resumed growth after some time (data not shown), explaining why the MIC is significantly higher than the acutely tolerated concentration. We determined the optimal stressor concentration for nisin to be 0.75 μg/ml (Fig. 1).



FIG 1 Effects of lantibiotic treatment on *B. subtilis* growth. *B. subtilis* was grown in defined medium to an  $OD_{500}$  of 0.35. Cultures were split and either treated with lantibiotics or left untreated. (A) Mersacidin, 30 µg/ml; (B) gallidermin, 6 µg/ml; and (C) nisin, 0.75 µg/ml. The time points of antibiotic addition are marked by arrowheads.

Characterization of the impact of lantibiotics on the B. subtilis cell wall. All tested lantibiotics were expected to influence cell wall integrity, as they prevent incorporation of cell wall precursors by binding lipid II. Their impact on cell wall integrity was investigated by microscopic examination of the cell shape after acetic acid-methanol fixation (Fig. 2, top) using a method described by Schneider et al., who investigated cell wall biosynthesis inhibitors vancomycin and plectasin (36). Upon inhibition of cell wall biosynthesis, small holes occur in the peptidoglycane layer, where new cell wall material is no longer incorporated. When the perforated cells are then fixed in a 1:3 mixture of acetic acid and methanol, the cytoplasmic membrane extrudes through holes in the cell wall, appearing as bubbles on the cell surface. To ensure that this phenomenon occurs specifically upon impairment of cell wall integrity and does not occur after treatment with agents impairing membrane integrity or simply as a result of sample handling, we used untreated, valinomycin-treated, and gramicidin A-treated B. subtilis cultures as negative controls. Valinomycin is a cyclic pep-



FIG 2 Influence of mersacidin, gallidermin, and nisin on *B. subtilis* cell wall integrity and membrane potential. (Top) Light microscopy images of *B. subtilis* fixed with acetic acid-methanol and immobilized with low-melting agarose show the influence of lantibiotics and comparator compounds on cell wall integrity. (Center) Fluorescence microscopy images of nonfixed cells show GFP-MinD localization after treatment with envelope-targeting agents. Cultures of *B. subtilis* 168 or *B. subtilis* 1981 GFP-MinD were grown until early exponential phase, divided into aliquots, and stressed with antibiotics for 15 min: (A) untreated control; (B) valinomycin (10  $\mu$ g/ml); (C) gramicidin A (0.025  $\mu$ g/ml); (D) gramicidin S (1  $\mu$ g/ml); (E) nisin (0.75  $\mu$ g/ml); (F) gallidermin (6  $\mu$ g/ml); (G) mersacidin (30  $\mu$ g/ml); (H) bacitracin (12  $\mu$ g/ml); (I) vancomycin (1.5  $\mu$ g/ml). (Bottom) Corresponding light microscopy pictures of the *B. subtilis* GFP-MinD cells shown above. Note that *B. subtilis* GFP-MinD mutants typically grow longer than wild-type *B. subtilis* due to MinD overexpression.

tide carrier ionophore that selectively transports potassium ions across the bacterial membrane, while peptide channel ionophore gramicidin A transports monovalent cations (18). Neither the untreated controls (Fig. 2A) nor the valinomycin or gramicidin Atreated cultures (Fig. 2B or C, respectively) displayed any membrane extrusions. As expected, all lantibiotics led to membrane extrusions (Fig. 2E, F, G), demonstrating their influence on cell wall biosynthesis in *B. subtilis*. Further, we tested the cyclic peptide antibiotic gramicidin S (Fig. 2D), which served as a positive control for depolarization experiments. Gramicidin S had been suggested to affect membrane-bound processes indirectly by causing delocalization of membrane-associated enzymes (25, 26, 44, 55). Indeed, after gramicidin S treatment, cells also showed membrane extrusions, corroborating an effect of gramicidin S on cell wall biosynthesis.

Lantibiotics' influence on B. subtilis membrane potential. In addition to cell wall biosynthesis inhibition, gallidermin and nisin are expected to influence membrane integrity due to integration into the membrane and pore formation, respectively. Therefore, we investigated the influence of lantibiotics on the membrane potential monitoring microscopically the localization of a GFP-MinD fusion protein (Fig. 2, central and bottom panels). MinD is part of the cell division regulation machinery of B. subtilis. As long as the membrane potential remains intact, MinD is attached to the membrane and localizes at the cell poles and in the cell division plain. It was shown that MinD localization is altered by membrane-depolarizing agents like valinomycin or proton ionophore carbonyl cyanide m-chlorophenylhydrazone (CCCP), due to the interruption of electrostatic interactions between MinD and the membrane surface (41). This phenomenon was employed here to investigate depolarizing properties of antibiotic agents. As described by Strahl and Hamoen (41), valinomycin was used as a positive control, as it causes rapid depolarization (Fig. 2B). Disruption of the membrane potential results in delocalization of GFP-MinD, which appears in fluorescing spots irregularly distributed throughout the cells. Nisin and gallidermin (Fig. 2E and F, respectively) exhibited the same GFP-MinD distribution pattern as valinomycin, indicating the collapse of the membrane potential (Fig. 2). In contrast, mersacidin did not disturb the polar and septal localization of GFP-MinD (Fig. 2G). These findings suggest that nisin and gallidermin but not mersacidin cause membrane depolarization in *B. subtilis*. Delocalization of GFP-MinD was also caused by the membrane-integrating antimicrobial peptide gramicidin S (Fig. 2D), which is neither an ionophore nor a pore former (25, 44).

Lantibiotics' ability to form pores in the *B. subtilis* membrane. In order to investigate the ability of mersacidin, gallidermin, and nisin to form pores in *B. subtilis*, we measured the lantibiotic-induced potassium release using a potassium-sensitive electrode (Fig. 3). Consistent with the literature, pore-forming lantibiotic nisin led to strong, immediate potassium release,



FIG 3 Potassium release from *B. subtilis* induced by mersacidin, gallidermin, nisin, and valinomycin. Potassium efflux from living cells was monitored with a potassium-sensitive electrode. Ion leakage is expressed relative to the total amount of potassium released after addition of the pore-forming lantibiotic nisin (100% value). Antibiotics were added at 50 s and applied at the same concentrations as for proteome analysis: 0.75 µg/ml nisin, 6 µg/ml gallidermin, 30 µg/ml mersacidin, and 10 µg/ml valinomycin. The symbol-free solid line represents the baseline without antibiotics.



FIG 4 Differential proteome analysis of *B. subtilis* 168 in response to mersacidin (A), gallidermin (B), and nisin (C). 2D gel-based protein synthesis profiles of the controls false colored in green were overlaid with those of the antibiotictreated samples false colored in red. In the overlays, downregulated proteins appear green, upregulated proteins appear red, and proteins expressed at equal rates appear yellow. Unidentified proteins are labeled as follows: M, induced by mersacidin; G, induced by gallidermin; N, induced by nisin; NGM, induced by nisin, gallidermin, and mersacidin.

whereas the potassium ionophore valinomycin (positive control) led to release of the same amount of potassium, following a slower kinetic characteristic for carrier ionophores. In contrast, as described previously, mersacidin did not cause an increase in extracellular potassium levels (6, 9, 14, 23). Gallidermin, which had not yet been tested regarding its ability to form pores in *B. subtilis*, caused only a minor potassium release of up to 20% compared to nisin, suggesting that it does not effectively form pores in *B. subtilis* under the tested conditions.

Proteomic response to lantibiotic treatment. Comparative proteome analyses were carried out to investigate whether or not the proteomic response would reveal marker proteins for the lantibiotic mechanism of action of cell wall biosynthesis inhibition by lipid II binding and how far it could provide insights into additional membrane-impairing properties of lantibiotics. To this end, B. subtilis was grown to early exponential phase and stressed with the appropriate concentrations of lantibiotics. [35S]-L-methionine pulse labeling was used to label selectively those proteins newly synthesized under antibiotic stress. This procedure allows monitoring of the acute bacterial stress response with exquisite sensitivity. The soluble intracellular protein fraction was then separated by 2D-PAGE. Representative proteome expression profiles of mersacidin, gallidermin, and nisin are shown in Fig. 4. Proteins induced at least 2-fold are referred to as marker proteins. They mirror the physiological stress conditions the cells are facing and proved valuable for studying antibiotic mechanisms of action.

Twenty-five (25) marker protein spots were induced by treatment with gallidermin, 13 by mersacidin, and 8 by nisin (Table 1). Using MALDI-TOF/TOF, we were able to identify 13, 7, and 6 protein spots, respectively. The remaining protein spots were clearly visible on autoradiographs reflecting protein synthesis after antibiotic treatment but did not accumulate to any appreciable extent. Although these unidentified protein spots do not allow rational insights into the antibiotic mechanism, they still serve as antibiotic-specific markers, as they are reproducibly induced after treatment.

### DISCUSSION

In this study, we set out to characterize the stress response of *B*. subtilis to three different lantibiotics with gradually different mechanisms of action regarding interactions with the cytoplasmic membrane but each inhibiting cell wall biosynthesis by binding to precursor lipid II. It was previously described that mersacidin acts solely on cell wall biosynthesis, whereas gallidermin is able to integrate into the membrane, although pores are formed only in some bacterial species and not others (8, 16). Nisin was shown to integrate into the membrane upon lipid II binding, forming pores of 2 to 2.5 nm in diameter (50), if concentrations exceed a certain threshold (8, 38). In preparation of the proteomic study, we tested nisin, gallidermin, and mersacidin for their effects on the B. subtilis cell envelope under growth conditions matching those of the proteomic profiling analysis. As expected, all lantibiotics significantly affected cell wall integrity as shown microscopically (Fig. 2). Protein localization studies using GFP-labeled MinD showed protein delocalization after treatment with nisin and gallidermin, indicating disruption of the membrane potential. Mersacidin tested negative for MinD-GFP delocalization. Significant potassium release demonstrating pore formation was observed only for nisin but not gallidermin or mersacidin (Fig. 2 and 3). Taken together, these results suggest that in *B. subtilis*, both nisin and gallidermin

TABLE 1 Responder pr	roteins induced	oy nisin, gallidermii	1, and mersacidin
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Responde protein       Nisin (25% inhibition)       Nisin (25% inhibition)       Gallidermin       Mersacian       Protein name       Function       Regulato         Lial H       6.6       19       8.6       9.2       Modulator of <i>lialHGFSR</i> operon expression       Function against cell envelope       Kell (2000)         YnrB       1.6       2.1       2.1       1.4       Similar to ABC transporter       Protection against cell envelope       SigW         NGMI       1.6       2.3       6.7       YnrB       Similar to ABC transporter       Stress       Energy limitation-mediated stress       SigB         YceC       1.6       2.3       6.7       YnrB       Similar to toxic anior resistance protein Stress response       SigW/Sig         YceC       1.3       4.2       Similar to toxic anior resistance protein Stress response       SigW/Sig         YceR       7.9       8.3       Not identified       Stress response       SigW/Sig         YufB       14       9.8       Similar to ABC transporter       Modification of Ipoteichoic acids       SigX         YufB       1.2       2.6       Pointhibio       Not identified       Stress response       SigX <td< th=""><th></th><th>Induction factor<sup>a</sup></th><th></th><th></th><th></th><th></th><th></th><th></th></td<>		Induction factor <sup>a</sup>							
LiaH6.6198692Modulator of lialHGFSR operon expression expressionTrmB3.42.89.18.6tRNA [guanin-N(7)]methyltransferase (RNA [guanin-N(7)]methyltransferaseYrrB1.62.12114Similar to ABC transporterNGMI1.62.5138.5Not identified Phage shock protein AProtection against cell envelope stressSigW stressNadE1.62.36.7NAD synthetaseEnergy limitation-mediated stress stress responseSigW/Sig SigW/SigYeeL134.2Similar to toxic anion resistance protein 14Stress responseSigW/Sig SigW/SigYrrB149.8Similar to ABC transporterStress responseSigW/Sig SigW/SigGM17.98.3Not identified p-3-Phosphoglycerate dehydrogenase Serine biosynthesis/oxidative stress responseStress responseSigXN1113Not identified PhopStress responseSigWYdB_n7UnknownStress responseSigWYdB_n72Poillin binding protein 3 RpSBCell wall biosynthesisSigWG33.9Not identified Stress responseSigWG42.4Not identified ResponseSigWG47.2Not identified 	Responder protein	Nisin (25% inhibition)	Nisin (50% inhibition)	Gallidermin	Mersacidin	Protein name	Function	Regulator	
TrmB     3.4     2.8     9.1     8.6     RKNA [guanin-N(7)]methyltransferase     Verter     1.6     2.1     21     14     Similar to ABC transporter       NGMI     1.6     2.5     1.3     8.5     Not identified     Protection against cell envelope     SigW       PspA     1.8     5.5     6.4     Phage shock protein A     Protection against cell envelope     SigB       NadE     1.6     2.3     6.7     NaD synthetase     Energy limitation-mediated stress     SigB       YceC     1.3     4.2     Similar to tellurium resistance protein     Stress response     SigW/Sig       YtrB     5.8     2.4     Similar to ABC transporter     Putative ABC transporte ATPase     SigW/Sig       GM1     7.9     8.3     Not identified     subunit     Subunit     Stress response     SigW/Sig       GM2     2.6	LiaH	6.6	19	86	92	Modulator of liaIHGFSR operon			
1rmb TrFE $3.4$ $2.8$ $9.1$ $8.6$ $tKNA [guann-M(7)]methyltransferaseNGM11.62.12114Similar to ABC transporterNGM11.62.5138.5Not identifiedPspA1.85.56.4Phage shock protein AProtection against cell envelopeSigWNadE1.62.36.7NAD synthetaseEnergy limitation-mediated stressSigBYceH5.82.4Similar to toki canior resistance proteinStress responseSigW/SigYreB149.8Similar to toki canior resistance proteinStress responseSigW/SigYreH149.8Similar to toki canior resistance proteinStress responseSigW/SigGM17.98.3Not identifiedsubunitsubunitGM27.98.3Not identifiedstress/NADH regenerationN11134.2D-Alanyiber-Alanine carrier proteinStress responseSigXYdBb7UnknownStress responseSigXYdBb7Pencilling protein 3Cell wall biosynthesisYdBb7Pencilling protein 52TranslationYdBb7Not identifiedIffeldGM37.2Not identifiedIffeldGM47.2Not identifiedIffeldGM57.2Not identifiedIffeldGM67.2Not identifiedIffeld$			• •			expression			
Yrrk1.62.12114Similar to ABC transporterNGM11.62.5138.5Not identifiedPspA1.85.56.4Phage shock protein AProtection against cell envelopeSigWNadE1.62.36.7NAD synthetaseEnergy limitation-mediated stressSigBYceC134.2Similar to tellurium resistance proteinStress responseSigW/SigYceH5.82.4Similar to toxic anion resistance proteinStress responseSigW/SigYtrB149.8Similar to ABC transporterPutative ABC transporterPutative ABC transporter ATPaseStress responseSigW/SigGM17.98.3Not identifiedsubunitVtrAStress responseSigWGM28.66.7Not identifiedstress/NADH regenerationVtrAStress response3.60.7Not identifiedstress responseSigXYdBa3.6UnknownStress responseSigXYdBa3.6UnknownStress responseSigXYdBa3.6UnknownStress responseSigXYdBa3.61.1Ribosomal protein 3Cell wall biosynthesisG17.2Not identifiedTranslationIf an additionG42.4Not identifiedIf an additionIf an additionG53.3Not identifiedIf an additionIf an additionG67.2Not identifiedIf addition <td>TrmB</td> <td>3.4</td> <td>2.8</td> <td>9.1</td> <td>8.6</td> <td>tRNA [guanin-N(7)]methyltransferase</td> <td></td> <td></td>	TrmB	3.4	2.8	9.1	8.6	tRNA [guanin-N(7)]methyltransferase			
NGMI PspA1.62.5138.5Not identifiedPspA1.85.56.4Phage shock protein AProtection against cell envelope stressSigWNadE1.62.36.7NAD synthetaseEnergy limitation-mediated stressSigBYceC134.2Similar to tellurium resistance protein stress responseStress responseSigW/SigYceH5.82.4Similar to ABC transporterPutative ABC transporter ATPase subunitYtrAGM17.98.3Not identifiedsubunitYtrAGM22.6 $-3$ -Phosphoglycerate dehydrogenase $-3$ -Phosphoglycerate dehydrogenaseSerine biosynthesis/oxidative stress responseSigXYdBb7UnknownStress responseSigXYvBb,7UnknownStress responseSigXYvBb,7UnknownStress responseSigXYvBb,7UnknownStress responseSigXYdBb3.6Not identifiedSigXSigW7UnknownStress responseSigXSigW7VolkientifiedSigXSigW3.6Not identifiedSigXSigW7Not identifiedSigXSigW7Not identifiedStress responseG17.2Not identifiedStress responseG23Not identifiedStress responseG43.9Not identifiedStressG53.3Not identified	YtrE	1.6	2.1	21	14	Similar to ABC transporter			
PspA   1.8   5.5   6.4   Phage shock protein A   Protection against cell envelope   Sigw     NadE   1.6   2.3   6.7   NAD synthetase   Energy limitation-mediated stress   SigB     YceC   13   4.2   Similar to tellurium resistance protein   Stress response   SigW/Sig     YceH   5.8   2.4   Similar to toxic anion resistance protein   Stress response   SigW/Sig     YtrB   14   9.8   Similar to ABC transporter   Putative ABC transporter ATPase   YtrA     GM1   7.9   8.3   Not identified   subunit   Stress response   SigW/Sig     GM2   2.6	NGM1	1.6	2.5	13	8.5	Not identified			
NadE1.62.36.7NAD synthetaseEnergy limitation-mediated stressSigBYceC134.2Similar to tellurium resistance proteinStress responseSigW/SigYceH5.82.4Similar to toxic anion resistance proteinStress responseSigW/SigYtrB149.8Similar to toxic anion resistance proteinStress responseSigW/SigGM17.98.3Not identifiedsubunitYtrAGM22.6Patative ABC transporterStress/NADH regenerationYtrAN1113Not identifiedStress/NADH regenerationSigWNIA113Not identifiedStress/NADH regenerationSigWYtBa3.6UnknownStress responseSigWYtBa7Penicillin binding protein 3Cell wall biosynthesisSigWYuBa7Penicillin binding protein 3Cell wall biosynthesisSigWYuBa3.9Not identifiedTranslationFerine biosynthesisG17.2Not identifiedFerine biosynthesisFerine biosynthesisG33.3Not identifiedFerine biosynthesisFerine biosynthesisG42.4Not identifiedFerine biosynthesisFerine biosynthesisG43.9Not identifiedFerine biosynthesisFerine biosynthesisG42.8Not identifiedFerine biosynthesisFerine biosynthesisG65.8Not identifiedFerine biosynthesisFerine bi	PspA	1.8	5.5	6.4		Phage shock protein A	Protection against cell envelope stress	SigW	
YccC134.2Similar to tellurium resistance protein Similar to toxic anion resistance protein Stress responseStress response SigW/Sig 	NadE	1.6	2.3	6.7		NAD synthetase	Energy limitation-mediated stress response	SigB	
YceH   5.8   2.4   Similar to toxic anion resistance protein Similar to ABC transporter   Stress response   SigW/Sig     GM1   7.9   8.3   Not identified   number   ytrA   subunit     GM2   8.6   6.7   Not identified   number   stress/NADH regeneration   ytrA     SerA   1.2   2.6   D-3-Phosphoglycerate dehydrogenase   Serine biosynthesis/oxidative stress/NADH regeneration   stress/NADH regeneration     N1   1   13   Not identified   stress response   SigW     YvIBa   3.6   Unknown   Stress response   SigW     YvIBb   7   Unknown   Stress response   SigW     YvIBa   7   Pencillin binding protein 3   Cell wall biosynthesis   SigW     YvIBa   7.2   Not identified   Translation   SigW     Stress response   5.9   Not identified   Stress response   SigW     GG1   7.2   Not identified   Stress response   SigW     GG3   3.9   Not identified   Stress response   SigW     GG4   2.4   Not identified   Stress response   SigW     GG5   3.3   Not identified   Stress response   SigW     GG4   <	YceC			13	4.2	Similar to tellurium resistance protein	Stress response	SigW/SigB	
YtrB   14   9.8   Similar to ABC transporter   Putative ÅBC transporter ATPase   YtrA     GM1   7.9   8.3   Not identified   subunit     GM2   8.6   6.7   Not identified     SerA   1.2   2.6   b-3-Phosphoglycerate dehydrogenase   Series bosynthesis/oxidative stress/NADH regeneration     N1   1   13   Not identified   stress/NADH regeneration     NtA   4.2   b-Alanylo-alanine carrier protein ligase   Modification of lipoteichoic acids   SigX     YvlB,   3.6   Unknown   Stress response   SigW     YvlB,   7   Unknown   Stress response   SigW     PbpC   27   Penicillin binding protein 3   Cell wall biosynthesis   SigW     G1   7.2   Not identified   Translation   Translation     G3   3.9   Not identified   G   G   G     G4   2.4   Not identified   G   G   G   G     G5   3.3   Not identified   G   G   G   G     G4   2.4   Not identified   G   G   G   G     G5   3.3   Not identified   G   G   G   G	YceH			5.8	2.4	Similar to toxic anion resistance protein	Stress response	SigW/SigB	
GM1   7.9   8.3   Not identified     GM2   8.6   6.7   Not identified     SerA   1.2   2.6   D-3-Phosphoglycerate dehydrogenase   Serine biosynthesis/oxidative stress/NADH regeneration     N1   1   13   Not identified   Modification of lipoteichoic acids   SigX     YvlBa   3.6   Unknown   Stress response   SigW     YvlBb   7   Unknown   Stress response   SigW     YvlBb   7   Unknown   Stress response   SigW     PppC   27   Penicillin binding protein 3   Cell wall biosynthesis   SigW     G1   7.2   Not identified   Translation   Tenslation     G2   3   Not identified   Tenslation   Tenslation     G3   3.9   Not identified   Tenslation   Tenslation     G4   2.4   Not identified   Tenslation   Tenslation     G5   3.3   Not identified   Tenslation   Tenslation     G6   47   Not identified   Tenslation   Tenslation     G4   2.8   Not identified   Tenslation   Tenslation     G4   2.8   Not identified   Tenslation   Tenslation  G6   47<	YtrB			14	9.8	Similar to ABC transporter	Putative ABC transporter ATPase subunit	YtrA	
GM2   8.6   6.7   Not identified     SerA   1.2   2.6   D-3-Phosphoglycerate dehydrogenase   Serine biosynthesis/oxidative stress/NADH regeneration     N1   1   13   Not identified     DltA   4.2   D-Alanyl:D-alanine carrier protein ligase   Modification of lipoteichoic acids   SigX     YvlBa   3.6   Unknown   Stress response   SigW     PbpC   27   Penicillin binding protein 3   Cell wall biosynthesis     RpsB   4.1   Ribosomal protein S2   Translation     G1   7.2   Not identified   Image: SigW     G2   3   Not identified   Image: SigW     G3   3.9   Not identified   Image: SigW     G4   2.4   Not identified   Image: SigW     G5   3.3   Not identified   Image: SigW     G6   47   Not identified   Image: SigW     G8   2.8   Not identified   Image: SigW     G9   11   Not identified   Image: SigW     YqiG   5   NADH-dependent flavin oxidoreductase   Electron transport     YqiG   7.2   Not identified   Image: SigW     YqiG   7.2   Not identified   Image: SigW	GM1			7.9	8.3	Not identified			
Serial 1.2   2.6   p-3-Phosphoglycerate dehydrogenase   Serine biosynthesis/oxidative stress/NADH regeneration     N1   1   13   Not identified   stress/NADH regeneration     DltA   4.2   p-Alanyl:p-alanine carrier protein ligase   Modification of lipoteichoic acids   SigX     YvlBa   3.6   Unknown   Stress response   SigW     YvlBb   7   Unknown   Stress response   SigW     PbpC   27   Penicillin binding protein 3   Cell wall biosynthesis     G1   7.2   Not identified   Translation     G2   3   Not identified   Ga     G3   3.9   Not identified   Ga     G4   2.4   Not identified   Ga     G5   3.3   Not identified   Ga     G6   47   Not identified   Ga     G8   2.8   Not identified   Ga     G9   11   Not identified   Ga     G4   2.8   Not identified   Ga     G9   11   Not identified   Mot identified     G4   7.3   Not identified   Mot identified     G4   2.8   Not identified   Mot identified     G9   11   Not identified	GM2			8.6	6.7	Not identified			
N1   1   13   Not identified     DtA   4.2   D-Alanyl:D-alanine carrier protein ligase   Modification of lipoteichoic acids   SigX     YdB <sub>a</sub> 3.6   Unknown   Stress response   SigW     YvlB <sub>b</sub> 7   Unknown   Stress response   SigW     PbpC   27   Penicillin binding protein 3   Cell wall biosynthesis   Tanslation     G1   7.2   Not identified   Tanslation   Tanslation     G2   3   Not identified   Tanslation   Tanslation     G3   3.9   Not identified   Tanslation   Tanslation     G4   2.4   Not identified   Tanslation   Tanslation     G5   3.3   Not identified   Tanslation   Tanslation     G6   47   Not identified   Tanslation   Tanslation     G7   5.8   Not identified   Tanslation   Tanslation     G7   5.8   Not identified   Tanslation   Tanslation     G7   5.8   Not identified   Tanslation   Tanslation     G9   11   Not identified   Tanslation   Tanslation     G9   11   Not identified   Tanslation   Tanslation     M1   7.3<	SerA	1.2	2.6			D-3-Phosphoglycerate dehydrogenase	Serine biosynthesis/oxidative stress/NADH regeneration		
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YvlBa   3.6   Unknown   Stress response   SigW     YvlBbb   7   Unknown   Stress response   SigW     PbpC   27   Penicillin binding protein 3   Cell wall biosynthesis     RpsB   4.1   Ribosomal protein S2   Translation     G1   7.2   Not identified   G     G2   3   Not identified   G     G3   3.9   Not identified   G     G4   2.4   Not identified   G     G5   3.3   Not identified   G     G6   47   Not identified   G     G7   5.8   Not identified   G     G8   2.8   Not identified   G     G9   11   Not identified   G     YqiG   5   NADH-dependent flavin oxidoreductase   Electron transport     M1   7.2   Not identified   M	DltA			4.2		D-Alanyl:D-alanine carrier protein ligase	Modification of lipoteichoic acids	SigX	
YvlBb     7     Unknown     Stress response       PbpC     27     Penicillin binding protein 3     Cell wall biosynthesis       RpsB     4.1     Ribosomal protein S2     Translation       G1     7.2     Not identified     G       G2     3     Not identified     G       G3     3.9     Not identified     G       G5     3.3     Not identified     G       G6     47     Not identified     G       G7     5.8     Not identified     G       G8     2.8     Not identified     G       G9     11     Not identified     G       G4     2.8     Not identified     G       G9     11     Not identified     G       G9     11     Not identified     G       M1     7.3     Not identified     G       M2     7.2     Not identified     M	YvlB.			3.6		Unknown	Stress response	SigW	
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G4   2.4   Not identified     G5   3.3   Not identified     G6   47   Not identified     G7   5.8   Not identified     G8   2.8   Not identified     G9   11   Not identified     YqiG   5   NADH-dependent flavin oxidoreductase   Electron transport     M1   7.3   Not identified     M2   7.2   Not identified	G3			3.9		Not identified			
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G6   47   Not identified     G7   5.8   Not identified     G8   2.8   Not identified     G9   11   Not identified     YqiG   5   NADH-dependent flavin oxidoreductase   Electron transport     M1   7.3   Not identified     M2   7.2   Not identified	G5			3.3		Not identified			
G7   5.8   Not identified     G8   2.8   Not identified     G9   11   Not identified     YqiG   5   NADH-dependent flavin oxidoreductase   Electron transport     M1   7.3   Not identified     M2   7.2   Not identified	G6			47		Not identified			
G8   2.8   Not identified     G9   11   Not identified     YqiG   5   NADH-dependent flavin oxidoreductase   Electron transport     M1   7.3   Not identified     M2   7.2   Not identified	G7			5.8		Not identified			
G9   11   Not identified     YqiG   5   NADH-dependent flavin oxidoreductase   Electron transport     M1   7.3   Not identified     M2   7.2   Not identified	G8			2.8		Not identified			
YqiG   5   NADH-dependent flavin oxidoreductase   Electron transport     M1   7.3   Not identified     M2   7.2   Not identified	G9			11		Not identified			
M1   7.3   Not identified     M2   7.2   Not identified	YqiG				5	NADH-dependent flavin oxidoreductase	Electron transport		
M2 7.2 Not identified	M1				7.3	Not identified	*		
M2 2.2 Natified	M2				7.2	Not identified			
INIS 3.2 INOT Identified	M3				3.2	Not identified			

<sup>a</sup> Bold numbers indicate reliable marker proteins, more than 2-fold induced in three independent biological replicates.

effectively depolarize the membrane, with only nisin forming pores. Our results indicate that the GFP-MinD delocalization observed for gallidermin may be due to membrane depolarization caused by the lantibiotic integrating into the membrane rather than to pore formation. As Christ et al. demonstrated, gallidermin leads to effective pore formation in lipid bilayers containing mainly acyl chains up to 15 C atoms in length. Limited pore formation is observed in membranes composed predominantly of 17 C lipid tails (16). Consistently, Bonelli et al. reported pore formation for Staphylococcus simulans and Micrococcus flavus (both 15 C) but not for Lactococcus lactis (17 C) (8). Additionally to the membrane thickness, a large amount of branched-chain fatty acids in the membrane seems to negatively affect pore formation by gallidermin (16). The B. subtilis membrane consists mostly of 15 C lipid chains but at the same time possesses about 80% branchedchain fatty acids (17). Therefore, it is likely that gallidermin is not able to form pores in this organism due to membrane density rather than thickness. However, integration of gallidermin into the membrane might affect the physicochemical properties of the membrane, resulting in depolarization and consequently MinD-GFP delocalization. Similar effects have already been described for cationic antimicrobial peptide gramicidin S, a depolarizing agent. At sublytical concentrations, gramicidin S seems to disturb the membrane structure leading to dysfunction of membrane protein complexes. It uncouples electron transport by inhibiting cytochrome bd quinol oxidase and consequently disrupts the electrochemical gradient (25, 30, 44, 55). Gramicidin S also affects cell wall integrity (Fig. 2), potentially also due to disturbing the membrane-bound cell wall biosynthesis machinery by integrating into the membrane. There are also reports that gramicidin S increases membrane permeability for potassium ions (26), which might be the effect rather than the cause of membrane depolarization (30).

Based on their common mechanism of inhibiting cell wall biosynthesis by binding lipid II, the three lantibiotics were ideally suited for comparative proteomic studies aimed at describing proteomic signatures for cell wall biosynthesis inhibition. At the same time, this collection of lantibiotics in combination with the existing library of proteomic response profiles could be used to correlate the differences in the proteome according to the differences in membrane interaction. Many of the identified marker proteins are known to be upregulated in response to cell envelope stress (Table 1). YvlB, for instance, belongs to the SigW regulon induced by cell envelope stress. Taking into account the lantibiotic proteome profiles as well as previously published reference patterns of an extensive antibiotic proteome response library (4), we delineated three groups of marker proteins (Table 2, Fig. 5): those indicative of cell wall biosynthesis inhibition, those indicating membrane stress, and those induced in response to general envelope stress.

Markers for interference with cell wall biosynthesis by lipid II binding. Four proteins, namely, LiaH, YtrE, and TrmB, as well as unidentified protein NGM1, were induced by all tested lantibiotics. LiaH belongs to the LiaRS regulon. It forms multimeric ringlike structures consisting of 36 identical subunits that protect cell envelope integrity probably by binding to the membrane upon

Compound	Mode of action	Cell wall integrity	MinD delocalization	Potassium leakage	NadE	PspA	YceC	YceH	TrmB	LiaH	YtrE	YtrB	NGM1
Valinomycin <sup>b</sup>	Carrier ionophore	_	+	+	х	х	х	x					
Gramicidin A <sup><i>a</i></sup>	Channel ionophore	_	+	+	х	х		x					
Gramicidin $S^b$	Membrane integration	+	+	_	x	x	x	x	x	x			
Nisin	Pore formation and cell wall biosynthesis	+	+	+	х	х			х	х	х		х
Gallidermin	Membrane integration and cell wall biosynthesis	+	+	-	х	х	х	х	х	х	х	х	х
Mersacidin	Cell wall biosynthesis	+	_	_			х	x	x	х	х	х	х
Bacitracin <sup>a</sup>	Cell wall biosynthesis	+	_	_			x			x	х	х	х
Vancomycin <sup>a</sup>	Cell wall biosynthesis	+	_	_			х				x	х	

TABLE 2 Marker proteins for antibiotics targeting membrane integrity and/or cell wall biosynthesis correlated with influence on cell envelope integrity<sup>c</sup>

 $\overline{a}$  Previously published (4).

<sup>b</sup> 2D gel images not shown.

<sup>c</sup> +, envelope functionality as measured in cell wall integrity assay, MinD delocalization assay, or potassium release assay was impaired; –, control-like envelope functionality; x, marker protein upregulated at least twofold in response to antibiotic treatment.

cell wall stress (51). LiaRS has been shown before to be induced by various cell wall biosynthesis inhibitors, such as bacitracin, nisin, ramoplanin, and to a lesser extent vancomycin, measuring *liaI*-*lacZ* reporter gene activation (29). Further, the LiaRS system was



FIG 5 Details of autoradiographs of 2D gels depicting the specific marker proteins for cell wall biosynthesis, membrane stress, and general cell envelope stress. Induction factors are displayed in the lower right corner as the average over three independent biological replicates.

induced after severe secretion stress, suggesting that protein accumulation in the cell envelope may also trigger LiaH induction (24). Besides the three lantibiotics investigated here, two inhibitors targeting membrane-bound steps of cell wall precursor synthesis were previously analyzed using proteomic profiling. Vancomycin prevents transglycosylation by binding to the amino acid side chain of lipid II, while bacitracin prevents recycling of the bactoprenol carrier that transports cell wall precursor molecules through the membrane (15). LiaH was upregulated in response to all lantibiotics, as well as bacitracin and gramicidin S, but not vancomycin. A possible explanation might be that the lipid II binding site of vancomycin is the amino acid chain of lipid II, which is remote from the membrane (15), while for all other compounds, the lipid II binding sites are in close proximity to the membrane.

YtrE and YtrB, which are upregulated by all investigated cell wall biosynthesis inhibitors with the exception of nisin, are encoded in the ytrABCDEF operon and form two cytoplasmic ATPase subunits of a putative ABC transporter (52). Earlier studies by Yoshida et al. demonstrated impaired acetoin import in ytr-deficient knockout mutants (53). However, it should be noted that the YtrF substrate recognition subunit shares homology with both antimicrobial peptide efflux systems and the FtsX permease involved in cell division. Delayed spore formation was observed in the absence of FtsX (21) as well as in ytr-deficient mutants (53), possibly indicating similar functions of both proteins. Although it is vet unclear how YtrE/YtrB induction might be connected to the antibiotic mechanism, they serve as specific markers for cell wall biosynthesis inhibition. YtrE is the only marker protein shared by all lantibiotics, vancomycin, and bacitracin. It is not induced by other antibiotics tested and is, therefore, the most reliable marker for inhibition of a membrane-bound step of cell wall biosynthesis.

TrmB is a tRNA-modifying [guanine-N(7)-]-methyltransferase involved in tRNA maturation (54). What role it might play in the response to lantibiotic treatment remains unclear. Like LiaH, unidentified NGM1 is induced by all lantibiotics and bacitracin but not vancomycin, demonstrating a different cell response to vancomycin.

Likewise, the response pattern induced by the lipopeptide friulimicin B differs from that of cell wall biosynthesis-targeting antibiotics that bind proximal to the membrane. In contrast to the lipid II-binding lantibiotics tested here, friulimicin B binds to bactoprenol monophosphate, thus inhibiting carrier recycling. As a result, cell wall biosynthesis and biosynthesis of wall teichoic acids and carbohydrate-containing capsular material are impaired, and cell envelope modification reactions such as glycosylations are inhibited. All these pathways depend on bactoprenol-mediated building block shuffling to the outside (37). As expected for an antibiotic interfering with cell wall biosynthesis, friulimicin B leads to membrane extrusions in the methanol-acetic acid assay. It does not depolarize the bacterial membrane in the GFP-MinD assay (data not shown) and does not cause potassium leakage (37). Friulimicin B shares a high mode of action similarity with bacitracin, which binds to bactroprenol pyrophosphate. However, while bacitracin induces all cell wall biosynthesis marker proteins, friulimicin B induces only the unidentified marker proteins NGM1 and GM1 but not LiaH or YtrE (data not shown). In a previous study, Wecke et al. further showed induction of YceC and YceH (46), which in the present study were upregulated in response to several of the cell envelope-targeting antibiotics but were not correlated strictly with either interference with the membrane or with cell wall biosynthesis.

A clear distinction based on the proteome response can be made between inhibition of membrane-bound steps of cell wall biosynthesis and inhibition of either intracellular or extracellular steps. This could be shown using D-cycloserine or methicillin, respectively. D-Cycloserine, which inhibits alanine racemase, and D-alanine:D-alanine ligase, two intracellular enzymes of the pathway (28, 43), elicit a completely different proteomic response with no marker proteins overlapping with any of the other investigated cell wall biosynthesis inhibitors (4). Methicillin inhibits penicillin-binding protein 2 (PBP-2), a membrane-anchored transpeptidase and transglycosylase (33); however, the methicillin binding site is not located near the membrane anchor, and a direct interaction of methicillin with the cytoplasmic membrane has not been reported. For this compound in B. subtilis, no acute stress response at all was observed on the proteome level (4), suggesting that B. subtilis cells do either not sense the consequences of methicillin action and/or are unable to react to them. Based on these results, differences in the proteomic response to cell wall biosynthesis inhibition relating to the localization of the steps inhibited and potentially even the proximity of the antibiotic binding sites to the bacterial membrane can be observed.

Marker proteins for interference with membrane integrity. PspA and NadE are both induced by nisin and gallidermin as well as gramicidin A, gramicidin S, and valinomycin. Among all agents tested so far, only compounds acting on the bacterial membrane led to induction of these two proteins, supporting that they are specific marker proteins for membrane stress.

Interestingly, upregulation of PspA and NadE was observed only at higher nisin concentrations, leading to approximately 50% growth inhibition, while the cell wall biosynthesis-specific markers LiaH and TrmB were upregulated already under moderate stress conditions, causing approximately 25% growth inhibition (Table 1). This is in line with a two-staged mechanism of action of nisin, which is characterized by lipid II complexation at lower concentrations and pore formation above a threshold concentration (9, 23). PspA, like LiaH, forms homomultimeric ringlike structures (22). Although Standar et al. showed that those 36-mers are able to form large scaffolds (40), more recent studies demonstrated localization of PspA at midcell and at the cell poles by use of a GFP fusion (19). It was shown that PspA is able to bind to membrane phospholipids and prevent proton leakage. It is thought to counteract external membrane damage and to stabilize the membrane during membrane transport (27). Structural homology between PspA and LiaH suggests similar functions in stabilizing the cell envelope by binding to the membrane surface. Our results indicate that PspA is responsive to membrane damage, while LiaH is induced by lipid II-mediated cell wall stress. However, the different triggers inducing both stress proteins remain to be identified (51).

NAD synthase NadE, which catalyzes the final step in NAD synthesis, is regulated on the transcriptional level by the house-keeping sigma factor SigA and by the alternative sigma factor SigB coordinating the general stress response in *B. subtilis* in response to environmental stress and energy limitation. As a consequence of antibiotic-related impairment of membrane integrity evidenced by the disturbed membrane potential, energy limitation is a possible explanation for NadE upregulation.

Differentiation between pore formation by nisin and membrane integration by gallidermin could not be securely established based on specific marker proteins. This may be due to rapid killing of cells once nisin concentrations reach the necessary threshold for pore formation, hardly leaving a chance for a coordinated stress response. For this reason, pore formation might be difficult to distinguish from other forms of membrane stress by proteome analysis alone. However, monitoring concentration-dependent killing kinetics already allows differentiation between compounds that integrate into the membrane and those that form pores. Rapid bacteriolytic pore formers leave only a very narrow concentration window for proteomic profiling experiments and should induce the membrane stress-specific marker proteins PspA and NadE at the appropriate antibiotic concentrations.

Marker proteins for envelope stress. Putative tellurium resistance protein YceC, induced by gallidermin and mersacidin, was also induced by several of the envelope-targeting compounds in our proteomic response library, namely, valinomycin, gramicidin S, bacitracin, and vancomycin, and can therefore serve as a general marker for envelope stress. Putative anion resistance protein YceH was also upregulated in response to gallidermin, mersacidin, valinomycin, gramicidin S, and gramicidin A. YceC and YceH are under the dual control of SigW and SigB, but to our knowledge their functions are yet unknown.

In conclusion, using lantibiotics and a number of other antibiotics with different targets related to the cellular envelope, we described marker proteins correlating with cell wall biosynthesis inhibition, membrane stress, and general cell envelope stress. The established marker proteins can now be used to characterize new antibiotic agents, particularly lantibiotics, with regard to their antibacterial mechanisms.

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